Connecting via Winsock to STN

```
Welcome to STN International! Enter x:x
```

LOGINID: SSSPTA1626KAS

PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

```
* * * * * * * *
                     Welcome to STN International
NEWS 1
                 Web Page URLs for STN Seminar Schedule - N. America
                 "Ask CAS" for self-help around the clock
NEWS
NEWS 3
         SEP 09 ACD predicted properties enhanced in REGISTRY/ZREGISTRY
NEWS 4
         OCT 03 MATHDI removed from STN
NEWS 5
         OCT 04 CA/Caplus-Canadian Intellectual Property Office (CIPO) added
                 to core patent offices
NEWS 6 OCT 13 New CAS Information Use Policies Effective October 17, 2005
NEWS 7 OCT 17 STN(R) AnaVist(TM), Version 1.01, allows the export/download
                 of CAplus documents for use in third-party analysis and
                 visualization tools
NEWS 8 OCT 27 Free KWIC format extended in full-text databases
NEWS 9 OCT 27 DIOGENES content streamlined
NEWS 10 OCT 27 EPFULL enhanced with additional content
NEWS 11 NOV 14 CA/Caplus - Expanded coverage of German academic research
NEWS 12 NOV 30 REGISTRY/ZREGISTRY on STN(R) enhanced with experimental
                 spectral property data
NEWS 13 DEC 05 CASREACT(R) - Over 10 million reactions available
NEWS 14 DEC 14 2006 MeSH terms loaded in MEDLINE/LMEDLINE
NEWS 15 DEC 14 2006 MeSH terms loaded for MEDLINE file segment of TOXCENTER
NEWS 16 DEC 14 CA/CAplus to be enhanced with updated IPC codes
NEWS 17 DEC 16 MARPATprev will be removed from STN on December 31, 2005
NEWS 18 DEC 21 IPC search and display fields enhanced in CA/CAplus with the
                IPC reform
NEWS 19 DEC 23 New IPC8 SEARCH, DISPLAY, and SELECT fields in USPATFULL/USPAT2
NEWS EXPRESS JANUARY 03 CURRENT VERSION FOR WINDOWS IS V8.01,
              CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP),
              AND CURRENT DISCOVER FILE IS DATED 19 DECEMBER 2005.
              V8.0 USERS CAN OBTAIN THE UPGRADE TO V8.01 AT
              http://download.cas.org/express/v8.0-Discover/
NEWS DCOST
              SINCE APPROXIMATELY 20:00 COLUMBUS TIME DECEMBER 29.
              SOME ONLINE COST DISPLAYS HAVE BEEN SHOWING COSTS IN
              2006 PRICES FOR STN COLUMBUS FILES. THIS HAS BEEN
              CORRECTED. PLEASE BE ASSURED THAT YOU WILL BE BILLED
              ACCORDING TO 2005 PRICES UNTIL JAN 1. PLEASE CONTACT
              YOUR LOCAL HELP DESK IF YOU HAVE ANY QUESTIONS. WE
              APOLOGIZE FOR THE ERROR.
```

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NEWS HOURS

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NEWS WWW CAS World Wide Web Site (general information)

Enter NEWS followed by the item number or name to see news on that specific topic.

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FILE 'HOME' ENTERED AT 07:20:30 ON 09 JAN 2006

=> file reg COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 07:20:41 ON 09 JAN 2006 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2006 American Chemical Society (ACS)

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STRUCTURE FILE UPDATES: 8 JAN 2006 HIGHEST RN 871465-69-9 DICTIONARY FILE UPDATES: 8 JAN 2006 HIGHEST RN 871465-69-9

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TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2005

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Structure search iteration limits have been increased. See HELP SLIMITS for details.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

Uploading C:\Program Files\Stnexp\Queries\10647191.str

Page 2 saeed

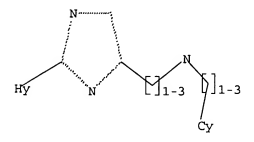
chain nodes :
6 7 8 9 10
ring nodes :
1 2 3 4 5
chain bonds :
2-6 5-7 7-8 8-9 9-10
ring bonds :
1-2 1-5 2-3 3-4 4-5
exact/norm bonds :
1-2 1-5 2-3 2-6 3-4 4-5 7-8 8-9 9-10
exact bonds :
5-7
isolated ring systems :
containing 1 :

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:Atom

L1 STRUCTURE UPLOADED

=> d L1 HAS NO ANSWERS L1 STR



Structure attributes must be viewed using STN Express query preparation.

=> s 11 SAMPLE SEARCH INITIATED 07:21:03 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 13407 TO ITERATE

14.9% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

1 ANSWERS

Page 3 saeed

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 261204 TO 275076
PROJECTED ANSWERS: 1 TO 289

L2 1 SEA SSS SAM L1

=> s 11 full

FULL SEARCH INITIATED 07:21:12 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 269257 TO ITERATE

100.0% PROCESSED 269257 ITERATIONS

258 ANSWERS

SEARCH TIME: 00.00.12

L3 258 SEA SSS FUL L1

=> file reg

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 170.46 170.67

FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 07:25:48 ON 09 JAN 2006 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2006 American Chemical Society (ACS)

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STRUCTURE FILE UPDATES: 8 JAN 2006 HIGHEST RN 871465-69-9 DICTIONARY FILE UPDATES: 8 JAN 2006 HIGHEST RN 871465-69-9

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH JULY 14, 2005

Please note that search-term pricing does apply when conducting SmartSELECT searches.

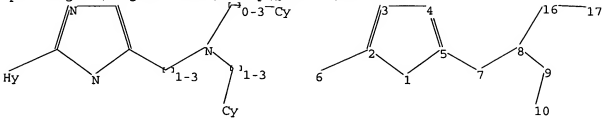
Structure search iteration limits have been increased. See HELP SLIMITS for details.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

=>

Uploading C:\Program Files\Stnexp\Queries\106471911.str



chain nodes :

6 7 8 9 10 16 17

ring nodes : 1 2 3 4 5 chain bonds :

2-6 5-7 7-8 8-9 8-16 9-10 16-17

ring bonds :

1-2 1-5 2-3 3-4 4-5

exact/norm bonds :

1-2 1-5 2-3 2-6 3-4 7-8 8-9 8-16 9-10 16-17

exact bonds :

4-5 5-7

isolated ring systems :

containing 1 :

Match level :

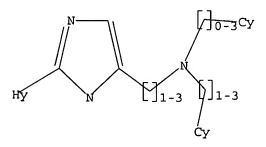
1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:Atom 16:CLASS 17:Atom

L4 STRUCTURE UPLOADED

=> d

L4 HAS NO ANSWERS

L4 `



STR

Structure attributes must be viewed using STN Express query preparation.

=> s 14

SAMPLE SEARCH INITIATED 07:26:14 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 13407 TO ITERATE

Page 5 saeed

14.9% PROCESSED 2000 ITERATIONS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

> BATCH **COMPLETE**

0 ANSWERS

261204 TO 275076 PROJECTED ITERATIONS:

0 TO PROJECTED ANSWERS:

0 SEA SSS SAM L4 L5

=> Uploading C:\Program Files\Stnexp\Queries\106471912.str

³0-3[−]Cy 17 Йу 10

chain nodes :

6 7 8 9 10 16 17

ring nodes : 1 2 3 4 5 chain bonds :

2-6 5-7 7-8 8-9 8-16 9-10 16-17

ring bonds :

1-2 1-5 2-3 3-4 4-5

exact/norm bonds :

1-2 1-5 2-3 2-6 3-4 4-5 7-8 8-9 8-16 9-10 16-17

exact bonds :

5-7

isolated ring systems :

containing 1 :

Match level :

1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:CLASS 8:CLASS 9:CLASS 10:Atom

16:CLASS 17:Atom

L6 STRUCTURE UPLOADED

=> d

L6 HAS NO ANSWERS

L6 STR

$$\begin{bmatrix} 1 \\ 0 \\ 3 \end{bmatrix} \begin{bmatrix} 1 \\ 3 \end{bmatrix}$$

Structure attributes must be viewed using STN Express query preparation.

0 ANSWERS

=> s 16 SAMPLE SEARCH INITIATED 07:27:20 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 13407 TO ITERATE

14.9% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**

PROJECTED ITERATIONS: 261204 TO 275076
PROJECTED ANSWERS: 0 TO 0

L7 0 SEA SSS SAM L6

=> s 16 full FULL SEARCH INITIATED 07:27:32 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 269257 TO ITERATE

100.0% PROCESSED 269257 ITERATIONS 16 ANSWERS SEARCH TIME: 00.00.12

L8 16 SEA SSS FUL L6

=> file caplus
COST IN U.S. DOLLARS
SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST
167.82 338.49

FILE 'CAPLUS' ENTERED AT 07:27:50 ON 09 JAN 2006
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Page 7 saeed

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FILE COVERS 1907 - 9 Jan 2006 VOL 144 ISS 3 FILE LAST UPDATED: 8 Jan 2006 (20060108/ED)

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http://www.cas.org/infopolicy.html

=> s 18

L9 6 L8

=> d ibib abs hitstr tot

L9 ANSWER 1 OF 6
ACCESSION NUMBER:
DOCUMENT NUMBER:
111LE:
INVENTOR(S):

INVENTOR(S):

PATENT ASSIGNEE(S):
SOURCE:

DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INSPONDATION:
FAILUT ACC. NUM. COUNT:
PATENT INSPONDATION:

CACCES COMMUNICATION
C

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. APPLICATION NO. KIND DATE DATE XIND DATE APPLICATION NO. DATE

A2 20041028 W 07 2004-US10626 20040408

A3 20050811

AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CR, CU, CZ, DE, DK, DM, DZ, EC, ER, EG, ES, FI, GB, GD, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LS, LT, LU, LY, MA, MD, MG, MK, MN, MW, MK, MZ, NA, NI, OM, FG, FH, FL, FT, KO, RU, SC, SD, SE, SG, SK, SL, SY, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, GH, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, ZW, AM, AZ, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, FR, GB, GR, HU, IE, IT, LU, MC, NL, FL, FT, RO, SE, SP, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, PATENT NO.

WO 2004091480

W: AE, AG,
CN, CO,
GE, GH,
LK, LR,
NO, NZ,
TJ, TM,
RW: BW, GH,
KS, FI,
SK, TR,
TD, TG
CA 2520255

EP 1611123

R: AT, BE, TD, TG
CA 2520255 AA 20041028 CA 2004-2520255 20040408
EP 1611123 A2 20060104 EP 2004-759191 2004008
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, PL, SK, HR
PRIORITY APPLN. INFO.: WANDER 141,382753 W0 2004-US10626 W 20040408

OTHER SOURCE(S): MARPAT 141:388753

AB The invention provides heterocyclic compds. for modulating protein kinase enzymic activity for modulating cellular activities such as proliferation, differentiation, programmed cell death, migration and chemoinvasion. Compds. of the invention inhibit, regulate and/or modulate kinases, particularly Tie-2. Methods of using the compds. and pharmaceutical compns. thereof to treat kinase-dependent diseases and conditions are also an aspect of the invention. Preparation of triazolyl compds. of the

invention is included. IT 783327-03-7

/84327-03-7
RE: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
(Biological study); USES (Uses)
(heterocyclic compound modulators of Tie-2 and other kinases, and
therapeutic use)
783327-03-7 CAPLUS
1-Naphthaleneacetamide, N-cyclopenty1-N-[2-[5-(4-methoxypheny1)-2-(4pyridiny1)-1H-imidazol-4-y1]ethy1]- (9CI) (CA INDEX NAME)

L9 ANSWER 2 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:327187 CAPLUS DOCUMENT NUMBER: 140:321364 Preparation of a control of a con 140:321364
Preparation of substituted imidazoles, pyrazoles and amides as high affinity C5a receptor modulators Thurkauf, Andrew: He, Xiao-shu; Zhao, He; Peterson, John; Zhang, Xiaoyan; Brodbeck, Robbin; Krause, James; Maynard, George; Hutchison, Alan Neurogen Corporation, USA U.S., 552 pp.
CODEN: USXXMM Patent
English
1 INVENTOR (S):

PATENT ASSIGNEE (S): SOURCE:

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
US 6723743	B1	20040420	US 2000-672071		20000928
<u>US 6884815</u>	B1	20050426	US 2003-461311		20030612
PRIORITY APPLN. INFO.:			US 1999-156390P	P	19990928
			US 2000-202749P	P	20000508
			US 2000-212499P	P	20000616
			US 2000-221787P	P	20000731
			US 2000-224036P	P	20000809
			US 2000-212449P	P	20000616
			US 2000-672071	A3	20000928

OTHER SOURCE(S): MARPAT 140:321364

AB The invention includes low mol. weight, non-peptidic, non-peptidommetic, organic mols. that can act as modulators of mammalian complement C5a receptors, preferably ones that act as high affinity C5a receptor ligands and also such ligands that can act as antagonists or inverse agonists of complement C5a receptors. Preferred compds. of the invention possess some or all of the following properties in that they are: (1) multi-aryl in structure: (2) heteroaryl in structure: (3) a pharmaceutically acceptable oral dose can provide a detectable in vivo effect; (4) comprise fewer than four or preferably no smide bonds, and (5) capable of habiting leukcoyte chemotaxis at nanomolar or sub-nanomolar concns. Such compds. include mainly substituted arylimidacoles! [m = 0-2; R] = H, OH, halo, NHZ, etc.; R2 = slkyl. cycloslkyl, haloalkyl, etc.; R31, R32, R5, R6 = H, OH, halo,

Page 9 saeed L9 ANSWER 1 OF 6 CAPLUS COPYRIGHT 2006 ACS OR STN

ANSWER 2 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)
NH2, etc.; R1 = alkyl, alkenyl, cycloalkyl, etc.; R7 = 0-3 groups selected from halo, NO2, CN, CF3, etc.], and also pyracoles, anides, etc. Betailed preps. of some of the title compds. was given. E.g., a multi-step synthesis of I (Arl = Ph; R1, R31, R32, R7 = H; R2 = Bu; R4 = 3,4-methylamedioxyphenyl) was presented. The invention also includes pharmaceutical compon. comprising the title compds. and the use of such compds. in treating a variety disorders.
439538-54-02 439536-56-22 439538-58-49
R1: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapoutic use); BIOL (Biological study); PREF (Preparation); USES (Uses)

(Uses)
(preparation of substituted imidazoles, pyrazoles and amides as high
affinity C5a receptor modulators)
439558-54-0 CAPLUS
HH-Inidazole-5-methanamine, N,N-bis(1,3-benzodioxol-5-ylmethyl)-1-butyl-2(2-thienyl)- (9CI) (CA INDEX NAME)

439558-56-2 CAPLUS
1H-Imidazole-5-methanamine, N-(1,3-benzodioxol-5-ylmethyl)-1-butyl-N-(phenylmethyl)-2-(2-thienyl)- (9CI) (CA INDEX NAME)

439558-58-4 CAPLUS

1H-Imidazole-5-methanamine, N-(1,3-benzodioxol-5-ylmethyl)-1-butyl-N-(cyclopentylmethyl)-2-(2-thienyl)- (9CI) (CA INDEX NAME)

ANSWER 2 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT 16

ANSWER 3 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN (Continued) cardiovascular, and immune system disorders. Pharmaceutical compns. contg. compds. of formula I are described. Addal., this invention provides labeled aminomethyl imidazoles compds., which are useful as probes for the localization of C5a receptors. Thus, II was prepd. from morpholine, bromodichlorobutylimidazole and (dihydrobenzodioxinylmethyl) (ethoxybenzyl)anine. Heny of the prepd. compds. exhibit a Ki value of less than 1 µM in an assay of C5a receptor mediated calcium mobilization. 656814-05-59 566814-05-69 666814-15-79
666814-05-59 566814-05-69 666814-15-79
666814-05-89 566814-15-79 (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
(preparation of aminomethyl imidazoles as complement C5a receptor L9 IT

(Uses)
(preparation of aminomethyl imidazoles as complement C5a receptor modulators)
66634-05-5 CAPLUS
1H-Imidazole-5-methanamine, 1-butyl-4-chloro-N-{(2,3-dihydro-1,4-benzodioxin-6-yl)methyl]-N-[(3-ethoxyphenyl)methyl]-2-(4-morpholinyl)-(SCI) (CA INDEX NAME)

666934-06-6 CAPLUS
IH-Imidazole-5-methanamine, 1-butyl-4-chloro-N-[(2,3-dihydro-1,4-bezodioxin-6-yl)methyl]-N-[(3-ethoxyphenyl)methyl]-2-(1-piperazinyl)-(9CI) (CA INDEX NAME)

666834-15-7 CAPLUS
Benzoic acid, 4-{[[[1-butyl-4-chloro-2-(4-morpholinyl)-1H-imidazol-5yl]methyl][[2,3-dhydro-1,4-benzodioxin-6-yl)methyl]amino]methyl]-, methyl
ester (9CI) (CA INDEX NAME)

L9 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2004:182872 CAPLUS
DOCUMENT NUMBER: 110:235712 CAPLUS
11TLE: Preparation of aminomethyl imidazoles as complement
C5a receptor modulators
Thurkauf, Andrew 2 hano, Her Zhang, Suoming; Gao, Yang
Neurogen Corporation, USA
PCT Int. Appl., 104 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent

Patent English

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE 20030821 A1 20040304 WO 2003-US26432 WO 2004018460

W: AE, AG, AL, M, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DX, PM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GH, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LK, LS, LT, LU, LY, HA, HD, MG, HK, MN, MY, KX, HX, NI, NO, NZ, OM, PG, PH, PL, PT, BO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZH, ZW

RW: GH, GM, KE, LS, MY, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TH, AT, BE, BG, CH, CY, CZ, DE, DX, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, HC, NL, PT, RO, SZ, SI, SK, TR, BF, BJ, CP, CG, CI, CM, GA, GN, GQ, GW, HL, HR, NE, SN, TD, TG

US 2004082577

A1 20040429 US 2003-647391 20030821

RIYY APPLN. INFO::

MARPAT 140:235712 WO 2004018460

PRIORITY APPLN. INFO.: OTHER SOURCE(S): GI MARPAT 140:235712

Aminomethyl imidazoles of formula I [R = H, halo, CN, alkyl, etc., Rl = H, OH, halo, amino, CN, nitro, alkyl, etc., R2 = alkyl, alkenyl, R3 = H, alkyl, R4 = alkyl, arylalkyl, etc., R5, R6 = H, alkyl, Ar = aryl, heteroaryl, fused Ph, etc., are prepared which are ligands of C5a receptors. Preferred compds. bind to C5a receptors with high affinity and exhibit neutral antagonist or inverse agonist activity at C5a receptors. The compds. can be used for the treatment of a variety of inflammatory,

ANSWER 3 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

666834-16-8 CAPLUS
Benzoic acid, 4-[{[[-butyl-4-chloro-2-{4-morpholinyl}-1H-imidazol-5-y|]methyl] (2,3-dihydro-1,4-benzodioxin-6-yl)methyl]amino]methyl]- (9CI) (CA INDEX NAME)

666834-22-6 CAPLUS
IH-Inidazole-5-methanamine, 1-butyl-4-chloro-N-[{2,3-dihydro-1,4-benzodioxin-6-yl)methyl]-N-[{3-ethoxyphenyl)methyl]-2-(1-piperidinyl)-{9CI} (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2006 ACS ON STN

ANSWER 4 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN (Continued) (phenylmethyl) -2-(2-thienyl)- (9CI) (CA INDEX NAME)

2

REFERENCE COUNT:

THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 4 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
111LE:
1203:818275 CAPLUS
139:286343
Combinantion therapy using a C5a antagonist and a C5a receptor-inactive therapeutic agent for the treatment of conditions with pathogenic inflammatory components
Krause, James
Neurogen Corporation, USA
PCT Int. Appl., 221 pp.
CODEN TYPE:
DOCUMENT TYPE:
DOCUMENT TYPE:
PATENT INFORMATION:
English
1 DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PRIORITY APPLN. INFO.: R SOURCE(S): MARPAT 139:286343 W 2003-U59424 W 20030327 R Compas. and methods for treating diseases that are associated with inflammation are provided. Such diseases include arthritis (particularly rheumatoid arthritis) and other autoimmune disorders, athma, cardio-and cerebrovascular disease, burns, psoriasis, reperfusion injury, and terumatic CNS and spinal cord injury. The compas, generally comprise at least one CSa antagonist and at least one CSa receptor-inactive therapeutic agent. The methods involve co-administration of at least one CSa antagonist and at least one CSa receptor-inactive therapeutic agent to a patient. The CSa antagonist and CSa receptor-inactive therapeutic agent to a patient. The CSa antagonist and CSa receptor-inactive therapeutic agent may be present within the same composition, or may be administered sep. to OTHER SOURCE(S): patient. 439558-56-2 īΤ 499558-56-7
RI: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
(Biological study); USES (Uses)
(C5s antagonist-C5s receptor-inactive therapeutic agent combination for treatment of condition with pathogenic inflammatory component)
439558-56-2 CAPLUS
HI-Imidzzole-5-methanamine, N-(1,3-benzodioxol-5-ylmethyl)-1-butyl-N-

L9 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER: 2003:796668 CAPLUS
DOCUMENT NUMBER: 139:307760
TITLE: Preparation of new aryl imid 139:307760
Preparation of new aryl imidazoles and related compounds as C5a receptor modulators Luke, George P., Haynard, George; Mitchell, Scott, Thurkauf, Andrew, Xie, Linghong; Zhang, Luyan; Zhang, Suoming; Zhao, He; Chenard, Bertrand L., Gao, Yang, Han, Bingsong; He, Xiao Shu Neurogen Corporation, USA PCT int. Appl., 356 pp. CODEN: PIXXD2 Patent INVENTOR (S): PATENT ASSIGNEE (S): SOURCE: DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. 20030228

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US 2004116424

Al 20040617

CA 2460888

AA 20031009

CA 2460888

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R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, ES, SI, LY, TY, LY, TY, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK

JF 2005528368

T2 20050922

US 2002-392145P

US 2002-392145P

US 2002-392145P

P 2002002626 PRIORITY APPLN. INFO.:

OTHER SOURCE(S):

L9 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

The title imidazoles, pyrazoles, pyridizines [1] the ring system in the formula I = 5-membered beteroaryl ring system (in which x = 0, A = C, N, O, S, and E and G = C, N, provided that the 5-membered heteroaryl ring system does not contain more than 3 heteroatems or more than 1 O or S stom) or 6-membered heteroaryl ring system (in which x = 1, A, B, E, and G = C, N, and provided that the 6-membered heteroaryl ring system does not contain more than 3 N atoms); N, R1 = H, OH, halo, stc., when E = N, then R2 = alkyl, alkenyl, CH2Ph, etc., when E = C, then R2 = H, halo, OH, etc., R3 = H, alkyl, alkenyl, etc., R4 = alkyl, alkenyl, cycloalkyl, etc., R5, R6 = H, alkyl, alkenyl, cycloalkyl, etc., R5, R6 = H, alkyl, z = 1-3, A1 = (un)substituted aryl, heteroaryl, Ph fused to 5-7 membered (un)saturated ring that has 0-2 ring atoms chosen from N, O, and S; Ar2 = cycloalkyl, cycloalkylakyl, aryl having 1 ring or 2 fused or pendant rings, etc., y = 1-6] which are ligands of C5s receptors, were prepared and formulated. E.g., s multi-step synthesis of II (starting from the benzimidate hydrochloride and 1-butylaniae), was given. Preferred compds. I bind to C5s receptors with high affinity (biol. data given) and swhibit neutral antagonist or inverse agonist activity at C5s receptors. This invention also relates to pharmaceutical compns. comprising such compds. It further relates to the use of such compds. in treating a variety of inflammatory and immune system disorders.
610287-35-89 & 100287-86-09 & 610285-41-59
610294-29-69 & 610295-02-69 & 610295-41-59
610294-29-69 & 610295-02-69 & 610295-41-59
610294-29-69 & 610295-02-69 & 610295-61-59
610294-29-69 & 610295-61-69 & 610295-61-610296-610-61005 & 610295 ΙT

6.1028-53-8P
RL: PAC (Pharmacological activity); SFN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
(preparation of new aryl imidazoles and related compds. as C5a receptor modulators)
610287-35-9 CAPLUS
1H-Imidazole-5-mechanamine, N,N-bis(1,3-benzodioxol-5-ylmethyl)-1-butyl-4-phenyl-2-(1H-pyrazol-1-yl)- (9CI) (CA INDEX NAME)

ANSWER 5 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

610295-02-8 CAPLUS
1H-Imidazole-5-methanamine, N,N-bis(1,3-benzodioxol-5-ylmethyl)-1-butyl-2-(3,5-dimethyl-1H-pyrazol-1-yl)-4-phenyl- (SCI) (CA INDEX NAME)

610295-41-5 CAPLUS lH-Imidazole-5-methanamine, 1-butyl-N-[{2,3-dihydro-1,4-benzodioxin-6-y|)methyl}-N-[{3-choxyphenyl}methyl]-2-{2-methyl-4-thiazolyl}-4-phenyl-(SCI) (CA INDEX NAME)

610298-53-8 CAPLUS
Benzamide, 4-[[[1-buty1-4-(4-methoxyphenyl)-2-(4-pyridinyl)-1H-imidazol-5yl]methyl](cyclohexylmethyl)amino]methyl]-2-hydroxy- (9CI) (CA INDEX
NAME)

L9 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

610287-86-0 CAPLUS
IH-Imidazole-5-methanamine, 1-butyl-N-{(2,3-dihydro-1,4-benzodioxin-6-yl)methyl}-N-[(3-ethoxyphenyl)methyl]-4-phenyl-2-(IH-pyrazol-1-yl)- (9CI) (CA INDEX NAME)

610288-15-8 CAPLUS
1H-Imidazole-5-methanamine, N-(1,3-benzodioxol-5-ylmethyl)-1-butyl-N-[{3-ethoxyphenyl)methyl}-4-phenyl-2-(1H-pyrazol-1-yl)- (9CI) (CA INDEX NAME)

610294-29-6 CAPLUS
1H-Imidazole-5-methanamine, 2-(1,3-benzodioxol-5-yl)-N,N-bis(1,3-benzodioxol-5-ylmethyl)-1-butyl-4-phenyl- (9CI) (CA INDEX NAME)

ANSWER 5 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

REFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L9 ANSWER 6 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN
ACCESSION NUMBER:
DOCUMENT NUMBER:
137:78952
ITITLE:
INVENTOR(5):
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DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

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	WO 2002049993 WO 2002049993						WO 2000-US26816						20000929				
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									,	WO 2	-0005	US 26	816		W 2	0000	929
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$$Ar^{1} \xrightarrow{N} \begin{array}{c} R^{1} \\ N \\ P_{2} \end{array} \xrightarrow{R^{3}} \begin{array}{c} R^{4} \\ Ar^{2} \end{array} \xrightarrow{N} \begin{array}{c} N \\ P_{2} \end{array} \xrightarrow{R^{3}} \begin{array}{c} R^{4} \\ Ar^{2} \end{array} \xrightarrow{N} \begin{array}{c} R^{4} \\ R^{4} \end{array}$$

The invention includes low mol. weight, non-peptidic, non-peptidommetic, organic mols. that can act as modulators of mammalian complement C5a receptors,

ANSWER 6 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN (Continued)

439558-58-4 CAPLUS
1H-Imidazole-5-methanamine, N-(1,3-benzodioxol-5-ylmethyl)-1-butyl-N-(cyclopentylmethyl)-2-(2-thienyl)- (9CI) (CA INDEX NAME)

ANSWER 6 OF 6 CAPLUS COPYRIGHT 2006 ACS on STN (Continued) preferably ones that act as high affinity C5s receptor ligands and also such ligands that can act as antagonists or inverse agonists of complement C5s receptors. Preferred compds. of the invention possess some or all of the following properties in that they are: (1) multi-aryl in structure; (2) beteroaryl in structure; (3) a pharmaceutically acceptable oral dose can provide a detectable in vivo effect; (4) comprise fewer than four or preferably no anide bonds, and (5) capable of habiting leukocyte chemotasis at nanomolar or sub-nanomolar concas. Such compds. include inidazoles I [R1 = H, OH, halo, etc.; R2 = alkyl, cycloalkyl, etc.; R3 + H, alkyl, atkenyl, cycloalkyl, etc.; R4 + alkyl, alkenyl, cycloalkyl, etc.; R5 + I, Ar2 = (un) substituted carbocyclic aryl, arylalkyl, etc., alkyl, alkenyl, cycloalkyl, etc.; Ar1 + (un) substituted carbocyclic aryl, arylalkyl, etc., etc. Datailed prepn. of some compds. I-III was given. E.g., a multi-step synthesis of I [Ar1 = Phr R1, R3 = H; R2 = Bur R4, R42 = 3,4-methylenedioxyphenyl) was presented. The invention also includes pharmaceutical compn. comprising such compds. I-III and the use of such compds. in treating a variety of inflammatory and immune system disorders.

Apsi8n=40-04 43958=56-24 439

(Uses)
(preparation of substituted imidazoles, pyrazoles and amides as high affinity C5a receptor modulators)
439558-54-0 CAPIUS
HH-Imidazole-5-methanamine, N,N-bis(1,3-benzodioxol-5-ylmethyl)-1-butyl-2-(2-thienyl)- (9CI) (CA INDEX NAME)

439558-56-2 CAPLUS
1H-Imidazole-5-methanamine, N-(1,3-benzodioxol-5-ylmethyl)-1-butyl-N-(phenylmethyl)-2-(2-thienyl)- (9CI) (CA INDEX NAME)

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